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Abstract: Multiphoton microscopy (MPM) has emerged as one of the most powerful and widespread technologies to monitor the activity of neuronal networks in awake, behaving animals over long periods of time. MPM development spanned across decades and crucially depended on the concurrent improvement of calcium indicators that report neuronal activity as well as surgical protocols, head fixation approaches, and innovations in optics and microscopy technology. Here we review the last decade of MPM development and highlight how in vivo imaging has matured and diversified, making it now possible to concurrently monitor thousands of neurons across connected brain areas or, alternatively, small local networks with sampling rates in the kilohertz range. This review includes different laser scanning approaches, such as multibeam technologies as well as recent developments to image deeper into neuronal tissues using new, long-wavelength laser sources. As future development will critically depend on our ability to resolve and discriminate individual neuronal spikes, we will also describe a simple framework that allows performing quantitative comparisons between the reviewed MPM instruments. Finally, we provide our own opinion on how the most recent MPM developments can be leveraged at scale to enable the next generation of discoveries in brain function.

DOI: <https://doi.org/10.1523/jneurosci.1527-18.2019>

Posted at the Zurich Open Repository and Archive, University of Zurich

ZORA URL: <https://doi.org/10.5167/uzh-184163>

Journal Article

Published Version



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Originally published at:

Lecoq, Jérôme; Orlova, Natalia; Grewe, Benjamin F (2019). Wide. Fast. Deep: Recent Advances in Multiphoton Microscopy of In Vivo Neuronal Activity. *Journal of Neuroscience*, 39(46):9042-9052.

DOI: <https://doi.org/10.1523/jneurosci.1527-18.2019>

Wide. Fast. Deep: Recent Advances in Multiphoton Microscopy of *In Vivo* Neuronal Activity

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Multiphoton microscopy (MPM) has emerged as one of the most powerful and widespread technologies to monitor the activity of neuronal networks in awake, behaving animals over long periods of time. MPM development spanned across decades and crucially depended on the concurrent improvement of calcium indicators that report neuronal activity as well as surgical protocols, head fixation approaches, and innovations in optics and microscopy technology. Here we review the last decade of MPM development and highlight how *in vivo* imaging has matured and diversified, making it now possible to concurrently monitor thousands of neurons across connected brain areas or, alternatively, small local networks with sampling rates in the kilohertz range. This review includes different laser scanning approaches, such as multibeam technologies as well as recent developments to image deeper into neuronal tissues using new, long-wavelength laser sources. As future development will critically depend on our ability to resolve and discriminate individual neuronal spikes, we will also describe a simple framework that allows performing quantitative comparisons between the reviewed MPM instruments. Finally, we provide our own opinion on how the most recent MPM developments can be leveraged at scale to enable the next generation of discoveries in brain function.

Key words: imaging; *in vivo*; multiphoton; two-photon

Introduction

When Adrian, Morison, and others performed the first *in vivo* recordings in anesthetized animals in the late 1930s, they were not able to record from more than a handful of neurons at a time (Adrian, 1936; Renshaw et al., 1940). Today, novel optical *in vivo* imaging methods enable routine high-resolution recordings of thousands of genetically identified individual neurons in the brain of awake, behaving animals over weeks and months (Trachtenberg et al., 2002; Holtmaat et al., 2009; Andermann et al., 2010; Attardo et al., 2015). Such capabilities illustrate the enormous advances that neurotechnologies have made over the last decades and how modern imaging tools open new avenues for neuroscience research. Compared with classical single-photon wide-field fluorescence microscopy, multiphoton microscopy (MPM) offers optical sectioning as well as depth penetration capabilities, two factors that greatly facilitate *in vivo* neural recordings deep in the intact living brain. Indeed, it was MPM that led the way for many important neuroscience discoveries revealing, among others, how new spines form during learning (Trachtenberg et al., 2002), how local spatial distribution of neuronal ac-

tivity patterns represents sensory information (Ohki et al., 2005), and the role of other types of brain cells, such as microglia (Nimmerjahn et al., 2005).

Since the first demonstration of a working two-photon (2P) microscope in 1990 (Denk et al., 1990), MPM technology has made remarkable progress, including the development of large FOV (Sofroniew et al., 2016; Stirman et al., 2016) and multibeam optical designs (Lecoq et al., 2014; Voigt et al., 2015), as well as high-speed laser scanning (Duemani Reddy et al., 2008; Grewe et al., 2010) and spatiotemporal multiplexing approaches (Cheng et al., 2011) (Fig. 1). Miniaturized, head-mounted versions of 2P microscopes were also developed (Helmchen et al., 2001; Piya-wattanametha et al., 2006) and have recently reemerged (Zong et al., 2017).

Although standard 2P microscopes are widely used in neuroscience, MPM still has some drawbacks and limitations in terms of the temporal resolution, large-scale network monitoring, long-term applicability, and its ability to record neural activity during natural behaviors in freely moving animals. In this review, we will mainly focus on the technological advances made to the initial MPM microscopy setup (for a basic schematic, see Fig. 2) over the past 10 years that, in our opinion, have the potential to substantially improve neuroscience research in the near future. Our goal is not to convey all relevant technical details but rather to make neuroscience researchers aware of recent technological developments in MPM as well as their basic strengths and limitations. Therefore, we organized this review to focus on developments for recording from a larger number of neurons, recording

Received March 2, 2019; revised Sept. 27, 2019; accepted Sept. 27, 2019.

We thank Kevin Takasaki and Peter Saggau (Allen Institute for Brain Science) for providing helpful comments on the manuscript; and Bénédicte Rossi for providing scientific illustrations. The authors thank the Allen Institute for Brain Science founders, Paul G. Allen and Jody Allen, for their vision, encouragement, and support.

The authors declare no competing financial interests.

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<https://doi.org/10.1523/JNEUROSCI.1527-18.2019>

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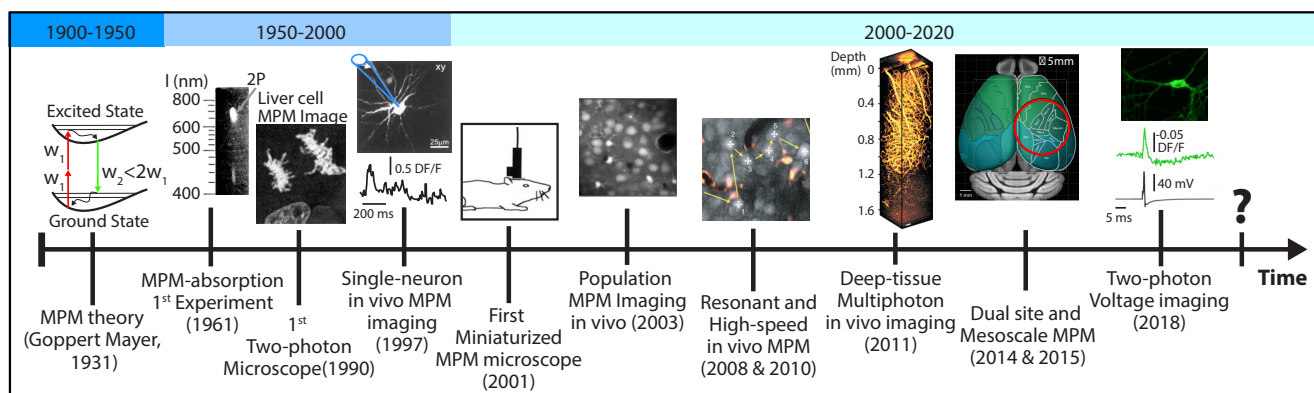


Figure 1. Historical timeline highlighting major milestones of MPM development and its applications. Adapted with permissions from Kaiser and Garrett (1961), Denk et al., (1990), Svoboda et al. (1997), Helmchen et al. (2001), Stosiek et al. (2003), Grewe et al. (2010), Sofroniew et al. (2016), Kobat et al. (2011), Chavarha et al. (2018).

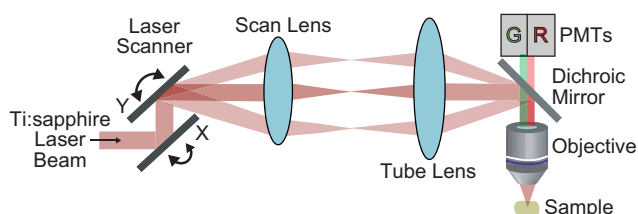


Figure 2. Schematic of a basic multiphoton microscope. An ultra-fast pulsed, near-infrared laser beam is directed onto a pair of X/Y scanning mirrors, which deflect the laser beam in two dimensions, depending on the voltage applied to their galvanometric elements. The beam then passes through the scan and tube lenses. The lens combination determines two fundamental parameters of scanning microscopy: the expansion of the beam entering the objective lens, which controls the size of the beam focus; and the scan angle of the laser beam into the objective, which is responsible for the size of FOV. Next, the laser beam is directed into the objective by a dichroic mirror that reflects near-infrared but passes light in the visible range. The objective lens focuses the short laser pulses into a tiny spot at the sample, resulting in multiphoton excitation of fluorescent indicator molecules. The emitted photons pass the dichroic mirror before being split into two channels (here, red and green) and are detected by two dedicated photomultiplier tubes (PMTs).

neurons at higher speed, and recording deeper in the brain; we also provide a comparison of MPM techniques. Due to space limitations, we do not cover any mobile and/or miniaturized MPM approaches that are described previously (Zong et al., 2017).

Meso-scale multiarea MPM approaches allow simultaneous *in vivo* recordings of thousands of neurons

Complex behavior often involves large brain networks that have both local and long-range connections and can span several millimeters in the mouse brain (Oh et al., 2014). To relate neuronal activity to behavior, it is crucial to simultaneously monitor the activity of such integrated long-range networks across many individual behavioral trials (Zhang et al., 2014). This experimental scientific need has fueled the development of new technologies that are capable of *in vivo* recordings from very large and distributed neuronal populations.

Interestingly, some of the biggest challenges in large-FOV optical systems were not optical. For example, only after the development of improved genetically encoded calcium indicators (GECIs) and related transgenic mouse lines (Dana et al., 2014; Madisen et al., 2015; Wekselblatt et al., 2016) did it become possible to label large populations of neuronal networks across multiple brain regions. In addition, early surgical protocols only

provided limited optical access to a single isolated, superficial brain region either by thinning of the animal's skull (Stosiek et al., 2003) or by implanting a small cover glass as skull replacement (Trachtenberg et al., 2002). Such limitations were overcome by protocols using large optical windows (i.e., 5 mm diameter) implanted over a large portion of the mouse dorsal cortex (Goldey et al., 2014). Recently, a novel glass implant preparation known as the “crystal skull” allowed imaging of the entire dorsal cortex by implanting a single, curved glass window that mimicked the bone plate curvature (Kim et al., 2016). Thanks to these innovations, imaging of large populations of neurons is no longer limited by methods of sample preparation and cell labeling; however, we still lack the imaging technology to sample all these neurons efficiently. To address this bottleneck, several optical MPM design approaches have been developed over the last decade. Broadly, these fall into two categories: single-beam scanning techniques, which sequentially traverse large fields of neural tissue at high speed; and multibeam scanning techniques, which can simultaneously interrogate different locations of the neuronal tissue.

Single-beam MPM for large-scale interrogation of brain tissue

Initial attempts to scale up neuronal recordings focused on optimizing single-beam scanning and extending MPM to the third dimension. By axially moving the microscope objective with a fast piezo actuator, smart 3D line-scanning approaches (Göbel et al., 2007) allowed recording from ~200 neurons at 10 Hz in the brain of live anesthetized rodents. An optimal use of the single laser-beam scanning time can be achieved by so-called random-access scanning (i.e., the laser beam is rapidly scanned over arbitrary selected points across the whole FOV). True random-access scanning can be realized with acousto-optic deflectors (AODs). These diffractive elements “deflect” a laser beam depending on the frequency of the sound wave generated by a piezo element that travels across a crystal positioned in the beam path. Two orthogonal AODs combine into a 2D scanner (Duemani Reddy et al., 2008; Grewe et al., 2010). If a pair of 2D-AODs are aligned in series with opposite sound propagation, 3D random-access imaging can be achieved with fast point-to-point transition times in the 10 μ s range (Katona et al., 2012; Nadella et al., 2016). This allows spending the entire laser beam scan time on neuronal somata while avoiding scanning any unlabeled neuropil. However, all line- and point-scanning techniques are sensitive to sample motion, which is typically present when imaging in awake animals (Dombeck et al., 2007; Nimmerjahn et al., 2009). Various approaches were proposed to mitigate this issue, for example,

by implementing a feedback loop for on-the-fly motion measurement and correction (Nadella et al., 2016) or by scanning local volumes around each area of interest (Szalay et al., 2016).

Despite these recent developments, fast raster scanning (i.e., sampling the FOV line by line) was favored for a long time, as motion artifacts could easily be resolved with custom postprocessing motion-correction algorithms (Kerr et al., 2005; Dombek et al., 2007). This encouraged widespread adoption of galvanometric scanners operating at their resonance frequency, enabling fast, 2D scanning rates at ~ 30 Hz (Fan et al., 1999). By combining such resonant scanners with an electrically tunable lens that shifts the axial focus of the laser beam by rapidly changing its shape, it became possible to scan two imaging planes at different depths at rates of up to 15 Hz, doubling the amount of recorded neurons (Grewe et al., 2011).

An alternative to inertia-limited axial scanners, such as piezo actuators or electrically tunable lenses, are spatial light modulators (SLMs) that are conjugated to the X/Y scanner and the Fourier plane of the objective (Liu et al., 2018). SLMs apply a variety of diffractive illumination patterns to the laser beam and thereby can shift MPM excitation laterally or axially. Contrary to inertia-limited electrically tunable lenses or piezo actuators, SLMs can provide fast (i.e., 5 ms) switching of the beam focus across the sample.

Another powerful concept for 3D imaging is remote focusing, which uses a second objective lens that focuses the beam on a mirror conjugated to the sample (Botcherby et al., 2012). Remote focusing not only corrects for optical aberrations introduced by the primary objective lens, but also provides a path for fast axial scanning as a lightweight mirror at the focus of the secondary objective can be moved by a fast actuator (e.g., a voice coil). Similarly, ultrasound lenses (Kong et al., 2015) refocus light by radially changing their refractive index. If used in a remote focusing setup, ultrasound lenses provide rapid axial scanning by changing the focus of the secondary objective.

An alternative approach to increase the number of recorded neurons is to enlarge the effective FOV by adjusting the microscope objective design. Commercially available objectives typically support scanning with a lateral extension of $500\text{--}800 \times 500\text{--}800 \mu\text{m}^2$, but they do not allow large scanning angles without deteriorating the optical resolution of the whole imaging system (Stosiek et al., 2003; Ji et al., 2016). This issue was resolved through tailored *in vivo* 2P imaging objectives for very large FOVs (Tsai et al., 2015; Sofroniew et al., 2016; Stirman et al., 2016). Combined with more sophisticated single-beam scanner arrangements, 1000–2000 neurons were recorded within a $5 \times 5 \text{ mm}^2$ FOV (Sofroniew et al., 2016) at a sampling rate of 10 Hz (Fig. 3A,B). However, large-FOV objectives with sufficient NA require large and heavyweight optics that cannot be moved quickly for fast axial scanning. Therefore, next-generation large-FOV systems rely on remote focusing, SLMs, or electrically tunable lenses (Sofroniew et al., 2016; Stirman et al., 2016). To further extend the FOV, custom postobjective scanners, which position the scanning beam in the sample space using pairs of micro-mirrors moved by a rotating stage, were recently introduced (Terada et al., 2018).

Despite the possibility that new *in vivo* preparations, such as the crystal skull (Kim et al., 2016), would now even allow recording from ~ 1 million cortical neurons, most of these single-beam scanning methods can only adequately detect calcium events across a very small subset of neurons simultaneously.

Multibeam MPM for sampling large neuronal population distributed across multiple distant sites

Perhaps one of the simplest multibeam imaging approaches is to use two independent pathways (Fig. 3C,D) by using micro-optics that provide the freedom of choosing two distant imaging sites (Lecoq et al., 2014). While this approach works well for distant recording sites ($>1\text{--}2$ mm apart), for adjacent areas, positioning multiple beams through a single objective can be more practical (Fig. 3E,F). For instance, with independent pathways, choosing new imaging areas requires a new optical alignment.

When designing multibeam scanning techniques, special attention must be given to crosstalk between the two excitation beams. Indeed, the presence of scattered photons in neural tissue (Oheim et al., 2001) limits the ability to later disambiguate multiple fluorescence emitters because fluorescence photons from one imaging plane are often detected in other imaging planes. This issue can be addressed either by *post hoc* source separation methods (Lecoq et al., 2014; Yang et al., 2016) or by temporal multiplexing schemes that support attributing the fluorescence signal to each individual scanning site (Cheng et al., 2011; Voigt et al., 2015). Temporal multiplexing can be easily implemented because MPM relies on high-power laser oscillators emitting very short (e.g., $\sim 100\text{--}200$ femtosecond laser pulses at 80 MHz repetition rate). These short laser pulses excite a fluorescence signal that lasts a couple of nanoseconds (GFP fluorescence lifetime ≈ 3 ns) (Striker et al., 1999). This allows the use of multiple laser excitation beams simultaneously but requires the pulses of each beamlet to be delayed in time. This delay is crucial because it later allows temporal separation of the individual fluorescence signals that have been excited by the different beams (Cheng et al., 2011; Voigt et al., 2015; Stirman et al., 2016).

Recently, reverberation optical loops that circulate (loop) the excitation beams several times through a single delay line were introduced to efficiently distribute laser pulses that differ in time and axial position (Beaulieu et al., 2018). Such temporal approaches typically allow multiplexing of up to 4 beams and thus increase the number of imaged neurons by a factor of 4 accordingly. However, introducing >4 beams would lead to an increased overlap of fluorescence decays in time and thus limit the ability to distinguish the signal sources. Another major drawback that might have limited the broad adoption of multiplexing MPM is the need for custom-built high-bandwidth electronics operating at least twice the rate of the multiplexed laser pulse rate. When multiplexing 4 beams from a standard 80 MHz laser, this results in a sampling frequency of > 640 MHz to comply with Nyquist sampling criterion.

An alternative to online demultiplexing are offline computational algorithms that remove multiplexing crosstalk (Mukamel et al., 2009; Pnevmatikakis et al., 2016), which, in combination with SLM-based multibeam microscope, allow *post hoc* source separation (Yang et al., 2016) (Fig. 3E,F). In this case, all simultaneously acquired planes are overlapping during the acquisition, and calibration data need to be collected during the experiment to estimate a (de)mixing matrix.

Custom beam engineering to match the excitation beam to fluorescently labeled neuronal populations

When dealing with sparsely labeled tissue, raster scanning techniques mostly sample nonlabeled pixels. One possible approach to minimize this issue is to elongate the point-spread-function (PSF) in the axial direction to more efficiently sample the tissue along this imaging axis. This so-called “Bessel beam scanning” projects a volume of tissue into a single plane (Lu et al., 2017) and

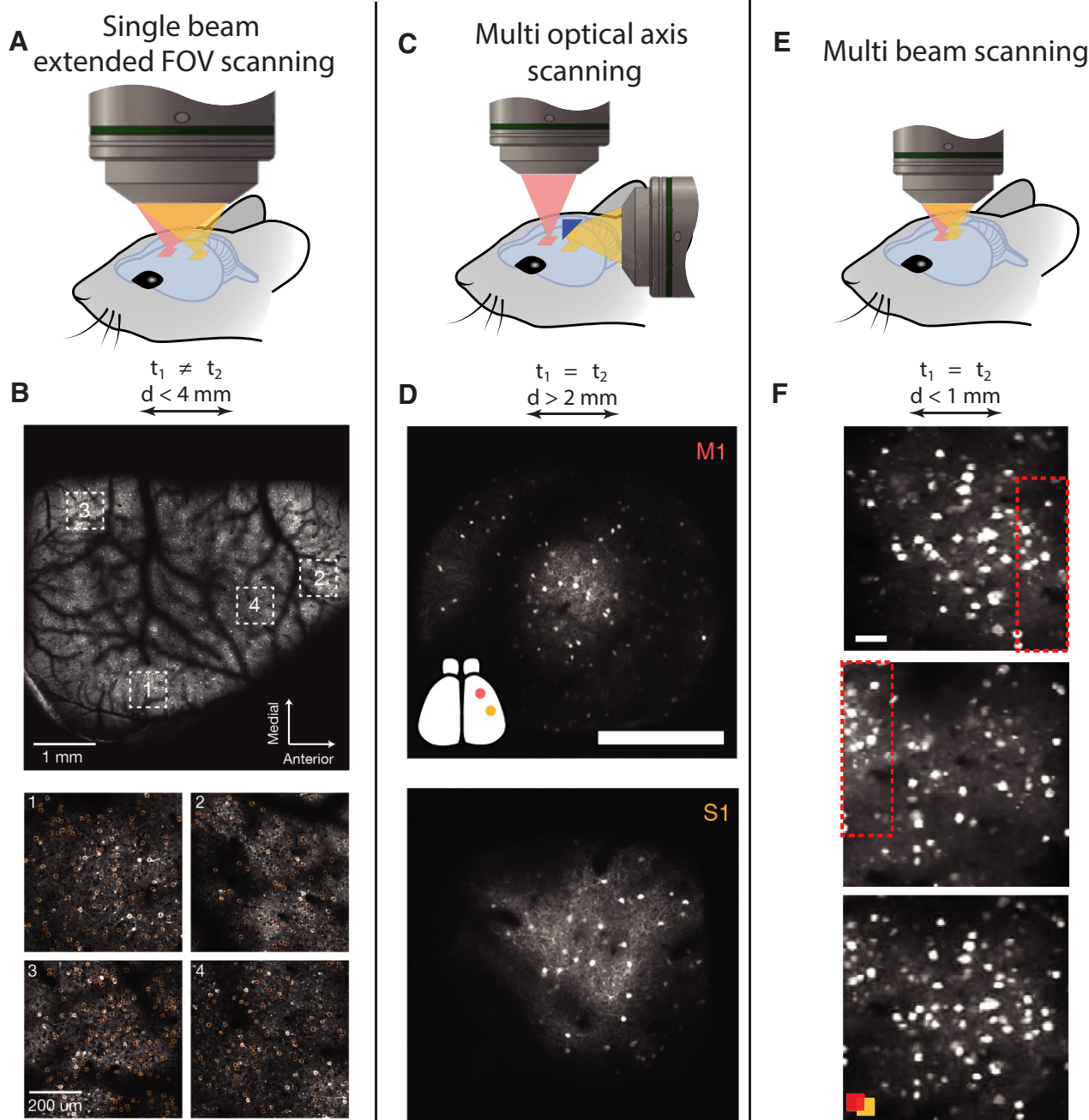


Figure 3. Large-area recording techniques. **A**, Schematic representation of a mesoscale (large FOV) imaging system known as 2P random-access microscope (2P-RAM). This imaging system supports slow scanning of a 5 mm^2 FOV or high-resolution imaging of multiple $500 \times 500 \mu\text{m}^2$ subregions with lateral separation of up to 4.5 mm. **B**, Low-magnification image acquired with the 2P-RAM system at 4.3 Hz frame rate, with an FOV of 5 mm^2 (top). Four separate areas of $500 \times 500 \mu\text{m}^2$ were sampled sequentially at 9.6 Hz (bottom). All data were acquired from a mouse expressing the calcium indicator GCaMP6f under a *thy-1* promoter. **C**, Schematic of the dual optical axis MPM system. Two excitation arms are fully independent and have separate optical paths. To solve geometrical constraints, one of the objectives is coupled with micro-optical elements positioned onto the mouse's head. **D**, Two distal areas (S1 and M1) were imaged with the dual-axis imaging system in a GCaMP6s mouse. Scale bar, $300 \mu\text{m}$. **E**, Schematic of a multibeam system with two simultaneous beams that scan two nearby areas. The scan areas were positioned within the FOV of a commercial objective and were laterally separated by $< 1 \text{ mm}$. **F**, Sequentially (top and middle) and simultaneously recorded images (bottom). Red boxes represent the area of FOV overlap. Computational algorithms and structural priors were used instead of excitation-emission encoding to separate the fluorescence signals of the two different planes. Scale bar, $50 \mu\text{m}$. Adapted with permission from Lecoq et al. (2014), Sofroniew et al. (2016), and Yang et al. (2016).

thus enables volumetric $400 \times 400 \times 100 \mu\text{m}^3$ MPM scan rates of 30–45 Hz. Although Bessel beams lack axial resolution along the z dimension, high-resolution image priors that are acquired with standard raster scanning can be used to retrieve the longitudinal coordinates of all recorded cells. Interestingly, Bessel beams at diameters well above the diffraction limit have also been successfully used in the context of densely labeled tissue. Indeed,

diffraction-limited PSFs are often unnecessary for *in vivo* imaging, and a loss in resolution can be traded off for higher imaging speeds (Lecoq et al., 2014; Stirman et al., 2016). In the extreme case, one can fully abandon the diffraction-limited PSF and very efficiently scan the tissue with a large PSF (e.g., $\sim 5 \times 5 \times 10 \mu\text{m}^3$) that approximately matches the neuronal soma size. Recent works from the Vaziri group (Prevedel et al., 2016; Weisen-

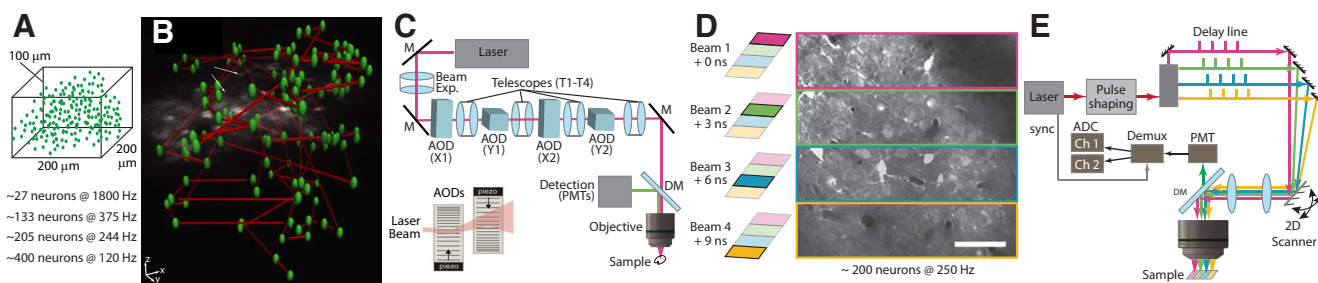


Figure 4. Techniques for large-volume and high-speed scanning using AOD scanners and spatiotemporal multiplexing. **A**, Schematic representation of 3D AOD scanning used in random-access multiphoton (RAMP) microscopy. In AOD MPM systems, the number of neurons within a 3D volume directly determines the temporal sampling rates. **B**, MPM image overlaid with one of the scan patterns from **A**. **C**, Schematic of RAMP microscope. A commercial $\text{Ti:Al}_2\text{O}_3$ laser beam is expanded to fill the AOD aperture. All AODs (X1...Y2) are optically conjugated via 4f telescopes (T1–T4). A dichroic mirror separates the IR excitation and visible light detection paths. GaAsP photomultiplier tubes (PMTs) detect emitted photons (for more technical details, see original publication, Cotton et al., 2013). **D**, Principle of spatiotemporal multiplexing where a larger FOV is simultaneously scanned with 4 beams to increase data acquisition rate by a factor of 4. **E**, Simplified layout of the microscope with spatiotemporal multiplexing. The laser emits pulses at 80 MHz corresponding to a 12.5 ns interpulse interval. The initial beam is divided into 4 beams, which are delayed by ~ 3 ns. All beams are recombined on a 2D-scanning module with different angular offset. In the imaging plane, this offset translates into a lateral displacement. Beam colors correspond to box colors in **D**. Emitted fluorescence is detected by PMTs, and photons are assigned to the correct image partition by a custom analog demultiplexing circuit based on their arrival time. For more details on temporal encoding and various techniques of multiplexing, see Cheng et al. (2011), Stirman et al. (2016), and Tsybolski et al. (2018). Adapted with permission from Cheng et al. (2011) and Cotton et al. (2013).

burger et al., 2019) explored this direction. By combining temporal focusing, beam engineering, and smart laser pulse energy control with fast laser scanning, they could record neuronal activity from up to $\sim 12,000$ neurons contained in a volume of $1 \times 1 \times 1.2 \text{ mm}^3$. This combination of methods, however, is technically challenging because it requires temporal focusing and a custom laser source to maintain sufficient signal-to-noise ratio (SNR) levels. In addition, larger PSFs effectively harm the ability to distinguish the activity of nearby cells, which has to be accounted for.

Fast MPM approaches to resolve neuronal activity at the millisecond time scale

Neuronal networks in the brain process incoming stimuli within a few tens of milliseconds. To understand such fast neuronal dynamics, it is of key importance to investigate neuronal activity patterns with an adequate temporal resolution. With the introduction of fast calcium indicators (Chen et al., 2013; Dana et al., 2019) and recent improvements of voltage indicators (Gong et al., 2015; Lin and Schnitzer, 2016; Kannan et al., 2018), high-speed imaging has become a major direction in MPM development (St Pierre, 2017; Chavarha et al., 2018). Strategies to realize fast imaging speeds *in vivo* share principles similar to those used for imaging large FOVs, but instead use the extra recording bandwidth to increase the sampling rate. Fast MPM approaches can be categorized into three classes that include the following: (1) single-beam line scanning methods or AOD-based scanning that allow fast random-access scanning of neuronal populations; (2) multibeam scanning techniques that divide the FOV into subregions; and (3) fast tomography-based systems that image the sample from multiple angles.

Early MPM approaches to improve imaging speed aimed to maximize the time that the laser focus is scanning structures of interest. Increasing the imaging speed to kilohertz rates became possible with AOD scanners that allow the laser beam to move within microseconds and hence enable true high-speed random-access imaging (Iyer et al., 2006). For example, a pair of AODs allowed 2D *in vivo* population calcium imaging with near millisecond precision (Grewe et al., 2010). Moreover, 3D AOD-based systems with fast axial focusing could image populations of neurons at kilohertz rates (Fig. 4A–C) (Katona et al., 2012; Cotton et al., 2013; Nadella et al., 2016). For these reasons,

AOD systems are particularly well suited for fast imaging of neuronal voltage signals. Recently, new genetically encoded voltage indicators have been combined with high-speed AOD-based MPM scanning approaches (Chavarha et al., 2018) to achieve fast population voltage imaging and to extract the spiking signals of individual neurons from the optical recordings with millisecond precision.

The second most common approach to increase imaging speed is to divide the FOV into several subfields and to use multiple beams that simultaneously scan these (Fig. 4D,E). Such parallelized raster scanning has been demonstrated using a fast resonant scanning system with four temporally multiplexed beams, resulting in scan rates of up to 250 Hz over $\sim 400 \mu\text{m}^2$ FOV (Cheng et al., 2011). Theoretically, many more beams can be used to perform *in vivo* imaging, even at kHz rates (US Patent 9,820,652). However, such systems would require proportionally more laser power to maintain similar SNR levels for each individual beam, and thus might quickly exceed the heat-absorption-caused damage threshold for neuronal tissues (Podgorski and Ranganathan, 2016). To further increase the imaging bandwidth, more sophisticated multiplexing schemes have been proposed that either rely on a digital micromirror device that encodes a specific digital signature to each pixel by flipping individual micromirrors of the array (Ducros et al., 2013) or using interferometric AOD schemes that encode each beam with a different beat frequency (Tsybolski et al., 2017). Although promising, these approaches have not yet been used extensively for *in vivo* recordings.

Another way to achieve kHz sampling rates relies on a tomography-like approach that scans the same probe from multiple angles (Kazemipour et al., 2019). These projections are acquired by four sequential line scans that traverse the FOV at different angles. This tomographic approach demonstrated dendritic imaging at rates >1 kHz across an FOV of $250 \times 250 \mu\text{m}^2$ at a depth of up to $250 \mu\text{m}$. However, one caveat of this approach is that the accuracy of signal reconstruction depends on the sparseness of the sample; for example, bright, clustered structures might not be reconstructed well. The tomography approach could be extended to larger FOVs; but as with other tomography methods, the number of reconstructed signal sources is limited to <1000 unless the number of angular projections is increased. Moreover, scanning with a complete

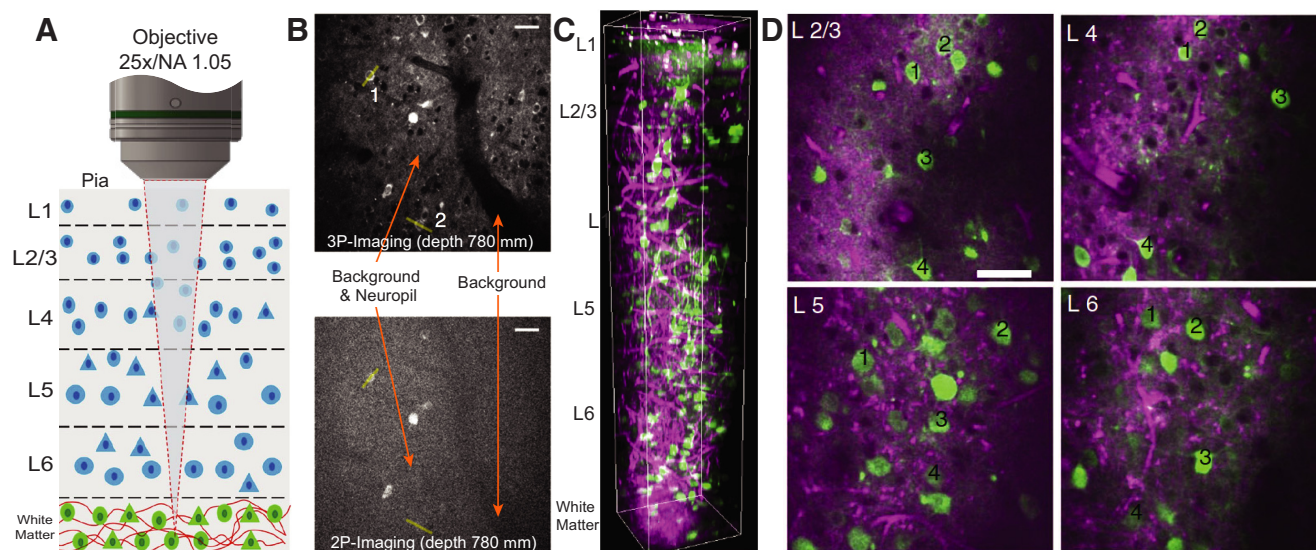


Figure 5. Advanced 3P microscopy approaches to record neuronal activity deep in the mouse brain. **A**, Schematic of the 3P deep-tissue imaging approach. **B**, Direct comparison of the signal-to-background ratio for 2P and 3P imaging of GCaMP6s-labeled neurons 780 μm deep in mouse cortex. “Background” arrow indicates the out-of-focus fluorescence measured inside a blood vessel. “Background & Neuropil” arrow indicates neuropil and the out-of-focus background. The two images have a comparable number of signal photons and are displayed with the same contrast settings (top 0.4% saturation). Scale bars, 30 μm . **C**, 3P z-stack acquired with 2 μm steps. Green represents GCaMP6s signal. Magenta represents the label-free third harmonic generation signal from the blood vessels in the visual cortex and myelin fibers in the white matter. Scale bar, 100 μm . **D**, Selection of 2D images from various layers (L2/3, L4, L5, and L6). Scale bar, 50 μm . FOV in all lateral images is 250 μm . Adapted with permission from Yildirim et al. (2019), and Ouzounov et al. (2017).

line also implies delivering much higher average power to the sample, which might lead to increased thermal load as well as photobleaching and toxicity.

MPM approaches to probe deep cortical and sub-cortical brain areas in awake animals

Measuring complex activity patterns that underlie behavior often requires recording neuronal populations in deep cortical or sub-cortical brain areas. However, imaging deeper than ~ 500 μm requires to adapt standard MPM techniques for an extended imaging depth. It is important to understand that the optical properties of tissue are the fundamental depth-limiting factors because both excitation and emission photons are highly scattered and absorbed by biological tissue. While the scattering problem could be addressed by increasing the laser intensity, this would introduce additional problems, such as sample heating, as well as out-of-focus and near-to-surface fluorescence volume excitation. Indeed, at least for 2P imaging, these issues seem most relevant for restricting the imaging depth (Theer and Denk, 2006), and the technologies that we discuss below attempt to resolve these limitations.

The most promising approach to increase the MPM imaging depth has been to shift the laser excitation light to longer wavelengths, for which scattering and absorption are substantially reduced (Lecoq and Schnitzer, 2011). Combined with the newest red-shifted GECIs, such as RCaMP, R-GECO, and jRCaMP (Akerboom et al., 2013; Dana et al., 2016), or related transgenic mice (Bethge et al., 2017), excitation wavelengths >1000 nm can be used to increase the 2P imaging depth down to ~ 0.8 mm (Ouzounov et al., 2017). While early approaches (Kobat et al., 2011) used optical parametric oscillators that provided an excitation wavelength of 1280 nm to image vascular structures down to 1.6 mm deep, functional (calcium) imaging beyond a depth of 1 mm (e.g., in mouse CA1) requires even longer wavelengths between 1600 and 1800 nm that still lie within the water absorption minimum. In this regimen, 3-photon (3P) absorption will become

the major process of fluorescence excitation. Indeed, out-of-focus and near-surface fluorescence volume excitation, the two biggest depth-limiting factors for 2P imaging, are substantially reduced for 3P imaging (Theer and Denk, 2006). Further increasing the laser excitation wavelength to 1700 nm can be achieved by another laser type, a photonic-crystal pumped by an erbium fiber laser. Using this approach, Horton et al. (2013) demonstrated 3P *in vivo* calcium imaging of neuronal and vascular structures up to a depth of 1.4 mm (Fig. 5). However, 3P imaging also requires an order-of-magnitude higher pulse energies to achieve the same fluorescence signal as 2P excitation. This is because fluorophore absorption cross-sections are substantially smaller for 3P excitation (compare 3P $\sim 10^{-83}$ $\text{cm}^6 \text{ s}^2/\text{photon}^2$ vs 2P $\sim 10^{-49}$ $\text{cm}^4 \text{ s}/\text{photon}$ for GFP) (Cheng et al., 2014). Such high pulse energies can be achieved through optical parametric amplifiers that provide pulse energies of hundreds of nanojoules within a tunable range of 1000–1600 nm. While the use of optical parametric amplifiers enabled structural *in vivo* imaging up to a depth of 1.5 mm at 1215 nm wavelength (Miller et al., 2017), more recently functional 3P imaging of mouse hippocampal and deep cortical areas at a depth of 0.7–1.2 mm has been demonstrated (Ouzounov et al., 2017; Yildirim et al., 2019) (Fig. 5).

To summarize, functional 3P microscopy is substantially more demanding than its structural counterpart as it requires fast scanning and short pixel dwell times to adequately sample the neuronal activity in time. In addition, pulse energies must be significantly higher (5–10 nJ compared with 0.3–3 nJ for standard 2P imaging) to gather enough signal during each pixel dwell time. Such high pulse energies also seem to increase photodamage in a nonlinear manner, which is more likely an issue in 3P imaging than heat absorption (Podgorski and Ranganathan, 2016; Yildirim et al., 2019). Another drawback of using optical parametric amplifiers for 3P imaging is that they can lower the pulse repetition rate to ~ 400 –800 kHz, which limits the number of pixels that can be scanned across the FOV within a certain time interval, thereby effectively reducing the FOV size. However,

Table 1. Representative publications for multiple recent advanced MPM techniques^a

Technique	No. of neurons, range	Recording frequency range, Hz	Depth, μm	Spatial resolution: lateral \times axial, μm	Reported application	Features and highlights	Reference
eMS2PM: encoded multisite 2PM	3–15	4100–1040	300	0.7×1.4	Voltage imaging in cultured cells, fast OGB calcium imaging <i>in vivo</i>	Simultaneous recording; less phototoxic; noise increase with increased number of recording sites; limited to 2D scanning; sensitive to motion	Ducros et al., 2013
RAPS: random-access pattern scanning	16–100	490–180	300	1.0×4.1	High SNR <i>in vivo</i> recordings of calcium transients	2D-AOD scanning; 1AP detection; sensitive to motion	Grew et al., 2010
SLAP: Tomographic TPM	100	1016	300	0.44×1.62	Single-cell voltage imaging, dendritic and axonal calcium recording	Constant frame rate independent of number of sources; tomographic image reconstruction; varying accuracy depending on label distribution; 2D FOV	Kazempour et al., 2019
ULOVE: 2D-AOD pattern scanning	1–150	15000–100	500	$0.5 \times 1.5^*$	Single-cell imaging of membrane activity in deep neurons during awake behavior	Resilience to motion; high SNR tracking of spikes; 2D FOV	Chavarha et al., 2018
Bessel 2PM: extended focus scanning	50–400	40–10	400	$0.5, 15–400$	High-speed volumetric calcium imaging <i>in vivo</i> in sparsely labeled tissue	Volumetric imaging with extended focus; volume size limited by power; axial dimension reconstructed <i>post hoc</i>	Lu et al., 2017
RAMP: 3D-AOD	27–411	1800–120	500	$0.5 \times 1.5^*$	High-speed calcium recordings from hundreds of neurons	Highest-speed sequential 3D recordings; sensitive to sample motion	Cotton et al., 2013; Nadella et al., 2016
Galvo-Galvo: raster scan with galvanometers	100–300	10–3	500	$0.5 \times 1.5^*$	Calcium imaging <i>in vivo</i>	Well-established technique; available from commercial companies	Denk et al., 1990
TPM-Mux: multibeam multiplexing	200–800	250–60	300	$0.5 \times 1.5^*$	Calcium imaging <i>in vivo</i> in multiple regions	Increased recording throughput by using multiple beams; crosstalk between beams	Stirman et al., 2016; Cheng et al., 2011
Dual-beam TPM: dual-area scan, two beams	200–600	10–3	700	1×9	Calcium imaging recordings <i>in vivo</i> in multiple regions	Simultaneous imaging of two areas decoupled in X/Y/Z; minimum separation is ~ 1 mm center-to-center; requires optical alignment for each experiment	Lecoq et al., 2014; Yang et al., 2016
s-Te-Flo: encoded multibeam scanning	200–4000	160–3	500	5×10	High-speed calcium recordings in populations of neurons	PSF sculpted to size of neuronal cell body-low spatial resolution; synchronizing pixels to laser pulses	Prevedel et al., 2016
Trepan2p: large FoV and beam multiplexing	400–5000	40–0.1	700	1.2×12	Dual beam calcium recordings over $>9 \text{ mm}^2$ FOV	Independent positioning of two beams within 9 mm^2 ; highly reconfigurable in terms of scan size and frame rate; crosstalk between beams	Stirman et al., 2016
2P-RAM: 25 mm^2 FOV	200–3000	44–1.9	500	1.2×5	Mesoscale imaging of calcium activity in neuronal populations	High collection efficiency; fast z scanning; highly flexible configuration of imaged regions and frame rates; requires prep adaptation to a bulky objective	Sofroniew et al., 2016
Resonant scanner, fast axial scanner	300–13000	30–2.5	500	$0.5 \times 1.5^*$	Calcium recordings <i>in vivo</i> in neuronal populations	Improved frame rate but lower SNR compared with Galvo-Galvo; available from commercial companies	Fan et al., 1999; Pachitariu et al., 2017

^aRecording capabilities and microscope parameters are listed along with the associated references.^{*}Spatial resolution values correspond to theoretical values for the objective lenses used in system designs, as PSF sizes were not reported.

newer 3P optical parametric amplifiers promise pulse repetition rates of up to 5 MHz. As for other technical developments, a combination of the techniques described above is currently explored to circumvent these limitations. For instance, combining Bessel beam with 3P absorption offers higher volume rate (Chen et al., 2018; Rodríguez et al., 2018).

Comparison of MPM techniques

The core idea of this review is to provide an overview of the most recent MPM developments and applications with the goal to en-

able researchers to select the best suitable method for their *in vivo* imaging experiments. However, a direct method comparison is difficult because comparative standards have not yet been established in the field. First, individual MPM approaches cannot be directly compared because experimental conditions vary across studies (e.g., fluorescent indicators, wavelengths, imaging depths, microscope objectives, photomultiplier tubes, and preamplifiers). Second, in addition to the sampling rate and number of neurons, key metrics needed to evaluate the quality of *in vivo* recordings, such as the ability to discriminate neuronal activity

signals from background, are often not reported. Third, typically only 1 or 2 performance data points (sampling rate and number of neurons) are reported per study, and the extrapolation to alternative MPM technologies proves difficult. Although a direct method comparison is hardly possible, it is clear that most recent MPM developments aimed to optimize the following three parameters: (1) the number of neurons that can be recorded; (2) the sampling rate and temporal precision to resolve neuronal activity; and (3) the reliability of detecting calcium events from the fluorescence signals and extracting individual neuronal spikes. While (1) and (2) are straightforward to evaluate, (3) is frequently underappreciated due to the lack of data. In the following section, we will attempt to analytically describe a framework to improve the standardization of reporting (3), the signal detection performance.

While simple metrics, such as the single action potential (1AP) SNR, were initially used to describe the ability to detect neuronal signals from fluorescence traces, recent modeling work suggests the use of more sophisticated metrics, which are commonly known in signal detection theory as the sensitivity index, or d' (Wilt et al., 2013). d' quantifies the ability to distinguish individual neuronal calcium events from baseline noise. Under certain assumptions (e.g., normal distribution), d' can be expressed as $d' = \frac{\mu_S^{\text{spike}} - \mu_N^{\text{no spike}}}{\sqrt{\frac{1}{2}(\sigma_S^2 + \sigma_N^2)}}$, where μ and σ are the mean and the SD

for the 1AP and the baseline signal distributions. Since σ is mainly driven by shot noise, one can assume $\sigma_N = \sigma_S$, which simplifies the equation to $d' = \frac{\mu_S^{\text{spike}} - \mu_N^{\text{no spike}}}{\sigma_S}$. As a rule of thumb, a d' of 5 relates to a correct 1AP detection rate of ~99%.

If the sampling rate f is substantially higher than the decay time constant τ_{ind} of the calcium indicator ($f \times \tau_{\text{ind}} \gg 1$), d' can be approximated as $d' \approx \frac{\Delta F}{F} * \sqrt{\frac{F_0 * \tau_{\text{ind}}}{2}}$ (Wilt et al., 2013).

Here, F_0 is the somatic background fluorescence in photons/s and $\Delta F/F$ is the peak fluorescence change detected for a single AP. This means that, to achieve a d' of 5 with GCaMP6f (1AP $\Delta F/F \sim 15\%$, $\tau_{\text{ind}} \sim 0.2$ s), the laser intensity or signal integration times need to be adjusted so that at least $F_0 > 11,000$ photons/s/cell are detected under baseline conditions. If the main interest is to reliably detect short bursts of, for example, 5 APs with the same d' , at least $F_0 > 2200$ photons/s/cell are required assuming a linear signal summation. In the following paragraph, we use standardized data acquired at the Allen Brain Observatory (de Vries et al., 2018) to provide an idea of how to calculate typical F_0 numbers and associated theoretical d' values for standard MPM systems.

The standard MPM systems at the Allen Institute for Brain Science use resonant galvanometric scanners, capable of scanning a single excitation laser beam across a $400 \times 400 \mu\text{m}^2$ FOV (512×512 pixels) at 30 Hz (de Vries et al., 2018). Taking into account the gain and offset of the data acquisition (photomultiplier tubes and preamplifiers), we calculated the median photon flux per pixel directly from the motion-corrected fluorescence data (for code to apply this method to any MPM data, see https://github.com/AllenInstitute/QC_2P) to be, on average, 0.89 photons per pixel per dwell time ($n = 2080$ experiments across both excitatory and inhibitory transgenic cre lines expressing GCaMP6f in all layers of the mouse visual cortex). In this case, a cellular soma spans ~ 100 pixels² that provide a $F_0 \equiv 2670$ pho-

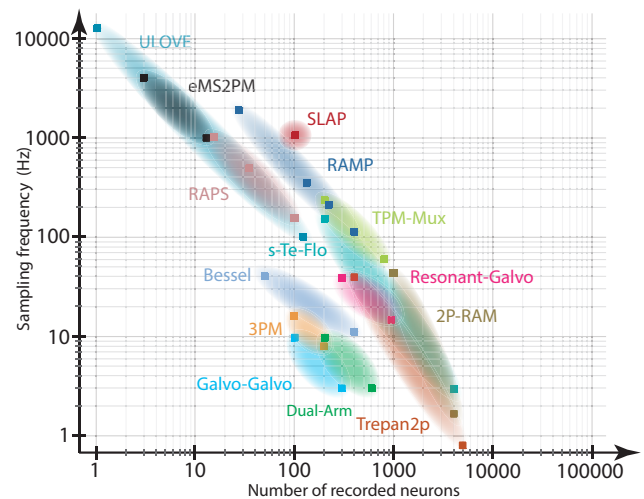


Figure 6. Comparison of MPM techniques in terms of number of neurons and recording speed. Mapping of recently developed MPM systems onto a 2D space (number of imaged neurons vs sampling rate). For some cases, we converted the imaging FOV into number of neurons assuming a densely labeled GCaMP mouse line, such as Slc17a7: Ai148 where, on average, 200 labeled neurons are detected in a $500 \times 500 \mu\text{m}^2$ FOV. Conventional galvo-galvo and resonant-galvo systems are included to emphasize the progress in technology development over the last decade. With the increasing number of scanning beams, the plot shifts toward higher speed or number of neurons, depending on the microscope configuration. Square markers represent published data. Clouds represent hypothetical interpolation between the reported data points.

tons/s/cell or a theoretical d' value of 2.4, which is well within a range where ≥ 5 APs are reliably detected ($d' \sim 5$).

A simple way to increase d' is to decrease the FOV size from $500 \times 500 \mu\text{m}^2$ to $180 \times 180 \mu\text{m}^2$, which would result in a $5\times$ increase of the number of pixels per neuron and thus a $5\times$ higher number of detected photons. This would result in $F_0 > 13,000$ photons/s/cell and thus a theoretical d' of 5.4. However, shrinking the FOV will also decrease the number of recorded neurons by a factor of 5. Alternatively, slower versions of GCaMP6 (i.e., GCaMP6s) can be used, providing an approximate 1AP $\Delta F/F$ change of 25% and a $\tau_{\text{ind}} \sim 0.6$ s (Chen et al., 2013). Because of the larger $\Delta F/F$ and τ_{ind} , GCaMP6s can achieve a d' of 5 with 1.7 times less recorded pixels per cell compared with GCaMP6f. However, using slower GECIs comes at the cost of a reduced temporal precision of detecting neuronal signals.

The best temporal precision can be achieved when using genetically encoded voltage indicators that directly report fast changes of the neuronal membrane voltage. For example, using an improved voltage indicator, ASAP3, *in vivo* population voltage imaging of 3–5 neurons was recently demonstrated at a rate of ~ 3 kHz, which is ~ 8 times faster than the temporal dynamics of the recorded voltage transients (Chavarha et al., 2018). The approximation introduced above (Wilt et al., 2013) will only hold if $f \times \tau_{\text{ind}} \gg 1$. Thus, if lower sampling frequencies are used, even higher photon counts are required to achieve a d' of 5. However, even when imaging at 3 kHz, imaging with a d' value of 5 can still falsely detect ~ 10 spikes per second, since each imaging frame can yield false positives. Thus, for fast voltage imaging, higher d' values are required. For example, sampling at 3 kHz, Chavarha et al. (2018) reported a d' of 9, which independent of the actual neuronal activity resulted in a false positive spike detection rate of one spike every 200 s and a false negative detection rate of 1.5×10^{-6} spikes per second. Given the reported 1AP $\Delta F/F$ of 9% and a decay time of ~ 4 ms, the number of baseline photons that

should be collected are $F_0 > 5000$ photons/ms/cell. These simple calculations illustrate that voltage imaging of large populations (>50 neurons) will require substantially more advanced MPM solutions or alternative imaging approaches that provide at least 1–2 orders of magnitude better imaging capabilities than current high-speed MPM approaches.

We summarize the properties of recent MPM developments in Table 1 and Figure 6, which shows that the number of recorded neurons is generally inversely related to the imaging rate. In this figure, we only report features and numbers that have been published or, in some cases, that we obtained by contacting the authors to clarify their method. Our comparison of the different microscopy systems is not optimal because the reported data have been acquired under different imaging conditions (e.g., differently labeling techniques, expression levels, and approaches to quantify and report SNR). The difficulty in comparing these methods also underlines the increasing need to develop a set of standardized evaluation metrics to report advances in MPM.

Discussion

The future of large-scale neuronal recording most certainly rests upon a combination of the above-mentioned techniques. Newly designed objectives lenses offer an extended view into neuronal tissues that can only be leveraged with high-bandwidth recording techniques. SLMs are promising tools providing flexibility to create multiple beams and even allow us to look deeper into tissue using concepts of adaptive optics (Ji et al., 2010; Liu et al., 2018). Similarly, multiplexing techniques combined with sophisticated scanning technologies, large FOV objective lens designs, and excitation beam shaping offer the opportunity to further increase the number of simultaneously recorded neurons to many thousands of neurons (Tsyboulski et al., 2018; Weisenburger et al., 2019). It is reasonable to expect that, in mice, entire cortical areas could be recorded in the very near future. As the sophistication of our instruments increases, the need for comparative quantitative standards becomes central to the development of future technologies. With integrated optical and mechanical computer-aided design tools, such as Zemax, Solidworks (Dassault Systèmes), and Creo (Parametric Technology), it becomes incredibly important to predict all performance aspects of new instruments before substantial financial investments are made.

Finally, given today's remarkable technological and experimental possibilities, the next generation of MPM instruments should be driven by neurobiological questions arising in the field. Indeed, multiphoton microscopes have matured from their initial design to showcase different, complementary capabilities. Instruments to image vast populations of neurons are designed very differently than those that aim to image dendritic processes.

Given the increased complexity of MPM instruments, their limited availability should not prevent us from concentrating on important neuroscience questions. In this regard, an academic observatory model currently used in high-energy physics and astronomy offers a unique opportunity to gain access to complex instrumentation and experimental techniques while encouraging data standardization. The idea of a brain observatory model is to open access to advanced experimentation based on unbiased scientific reviews of submitted proposals. Supportive staff can then perform the experiments while meeting high-quality standards and standardized operating procedures. Such an observatory approach is currently explored for systems neuroscience at the Allen Institute for Brain Science through the OpenScope project.

While monitoring the activity of thousands of neurons chronically was a distant vision just a few decades ago, it is now rou-

tinely performed in many laboratories through MPM of genetically encoded calcium reporters of neuronal activity. Significant technological developments have recently broadened our access to neuronal tissues and greatly increased the number of different techniques available. These allow for sampling larger populations of neurons, accessing neuronal networks located deep in the neuronal tissue, or achieving faster recording rates. Future developments will likely combine these techniques to further broaden our access to neuronal activity. Tight integration of these developments with neuroscience research is bound to transform our understanding of brain function.

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